Docket No.: 2870-0264P

**REMARKS** 

Claims 1-5, 10-16 and 18-21 are pending.

Claims 6-9 and 17 have been canceled without prejudice to filing the subject matter of

these claims in a divisional application.

Claim 12 has been amended to recite the subject matter of canceled claim 17.

No new matter has been added by way of the above-amendment.

Miura et al, U.S. Patent No. 6,248,512

Claims 1-5, 10 and 11 are rejected under 35 U.S.C. 102(e) and 103(a) as being

unpatantable over Miura et al. Applicants respectfully traverse these rejections.

Applicants have reviewed the Examiner's comments in the outstanding Office Action and

find that the Examiner has maintained this rejection based on a misunderstanding of the scope of

the compounds encompassed by Formula (IV) of instant claim 1. It is Applicants' position that

Miura et al. fail to teach or fairly suggest the compounds of Formula (IV) of instant claim 1.

Furthermore, it is Applicants' position that the compounds of Miura et al. do not satisfy the

requirement of instant claim 1 that the compounds have a hydrogen bond formation rate constant

 $K_{\Gamma}$  of 20-4000.

The Examiner alleges on page 3 of the outstanding Office Action that compounds A-29

and A-36 of Miura et al. fall within the scope of Formula (IV) of instant claim 1. This is

absolutely wrong. Compounds A-29 and A-36 are as follows:

MSW/GMD/mua

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The Examiner will note that Compounds A-29 and A-36 of Miura et al. require that a halogen is bonded to the nitrogen in the amide group.

This is in distinction to instant formula (IV), wherein the groups of R<sup>41</sup> and R<sup>42</sup> which bond to the nitrogen are not halogen. When R<sup>43</sup> is -N(R<sup>44</sup>)(R<sup>45</sup>), R<sup>44</sup> and R<sup>45</sup> are not halogen. Thus, the structure of Miura et al. is represented by -C(=O)-N-halogen whereas the structure of the formula (IV) is represented by -C(=O)-N-(alkyl, aryl or heterocyclic group). Accordingly, Compounds A-29 and A-36 are not encompassed by the present claims.

Furthermore, Compounds A-29 and A-36 do not satisfy the requirement of instant claim 1 that the compounds have a hydrogen bond formation rate constant  $K_f$  of 20-4000. This is explained in further detail below.

The Examiner further alleges on page 3 of the outstanding Office Action that compounds B-1, B-2, B-3, B-5, B-9 to 12 of Miura et al, also fall within the scope of Claim 1 in the present application. This is also wrong.

In addition, none of compounds B-1, B-2, B-3, B-5, B-9 to 12 nor compounds A-29 and A-36 satisfies the K<sub>f</sub> condition of Claim 1.

Kf is the hydrogen bond formation rate constant and is used as an index of hydrogen bond formation. It is fully explained in R.W. Taft et al., *J. Am. Chem. Soc.*, 91, 4794-4800 (1969), a copy of which is enclosed for the Examiner's review. (See the paragraph bridging pages 13 and 14 of the specification of the present application.)

Kf is the equilibrium constant in a reaction where a hydrogen bond is formed between p-FC<sub>6</sub>H<sub>4</sub>OH and a compound as follows:

The Examiner will note that each of Compounds A-29, A-36, B-1, B-2, B-3, B-5, B-9 to 12 of Miura et al. have an amide group. A hydrogen bond is generally formed between the oxygen atom of the carbonyl group in the amide and the hydrogen atom of the hydroxy group in p-FC<sub>6</sub>H<sub>4</sub>OH.

With respect to Compounds A-29 and A-36, both the amide groups are similar in that a bromine atom is bonded directly to the nitrogen atom. For example, the following hydrogen bond is formed between compound A-29 and p-FC<sub>6</sub>H<sub>4</sub>OH:

The oxygen atom of the carbonyl group has an acceptability of the hydrogen atom which depends on electron density on the oxygen atom. Generally, the lone electron pair on the right nitrogen atom of the above amide group can be shifted toward the oxygen atom by a resonance effect. Please however note that a bromine atom is bonded to the nitrogen atom in compound A-

29. It is well known in the art that a bromine atom withdraws electrons strongly. This results in the lone electron pair on the nitrogen atom in compound A-29 being strongly withdrawn to the bromine atom. The lone electron pair is not shifted toward the oxygen atom of the carbonyl group and therefore the oxygen atom has substantially no acceptability of the hydrogen atom. The case is the same as compound A-36. These facts are easily recognized by a person skilled in the art.

With respect to Compounds B-1, B-2, B-3, B-5, B-9 to 12, the Examiner will note that each compound actually has two amide groups along with HBr and Br-Br. For example, Compound B1 would have the following configuration wherein HBr forms hydrogen bond with the amide group as follows:

B-1

A hydrogen bond is formed between the oxygen atom of the carbonyl group and the hydrogen atom of HBr and therefore the oxygen atom has substantially no acceptability of the hydrogen atom of p-FC<sub>6</sub>H<sub>4</sub>OH. Compounds B-1, B-2, B-3, B-5, B-9 to 12 each has two amide groups in a molecule. One amide group forms a hydrogen bond with HBr and the other amide group forms a hydrogen bond with Br-Br. The oxygen atom of the other amide group also has substantially no acceptability of the hydrogen atom of p-FC<sub>6</sub>H<sub>4</sub>OH. The case is the same as compounds B-2, B-3, B-5, B-9 to 12. These facts are easily recognized by a person skilled in the art.

Thus, compounds A-29, A-36, B-1, B-2, B-3, B-5, B-9 to 12 do not satisfy the Kf condition of Claim 1.

Since Miura et al. only disclose compounds (even generically) having the nitrogen 10 MSW/GMD/mua

bonded to a halogen atom, Miura et al. do not anticipate nor render obvious the claimed invention. Thus, the claimed invention is patentable over Miura et al. and withdrawal of the rejection is respectfully requested.

## Bojora et al. U.S. Patent No. 3,667,959 and Miura et al.

Claims 12-21 are rejected under 35 U.S.C. 103(a) as being unpatentable over the combination of Bojora et al. U.S. Patent No. 3,667,959 and Miura et al. Applicants respectfully traverse the rejection.

First, Bojora et al. and Miura et al. are silent of polyhalogenated compounds. No one skilled in the art would have been motivated to use the polyhalogenated compounds in a photothermographic material and would have predicted the advantageous effects caused by utilizing the polyhalogenated compounds before the claimed invention was made.

The Examiner states in Item 5 of the official action that Bojora et al. disclose a photosensitive and thermosensitive element containing sulfonyl group including that claimed in the present claimed invention in a photothermographic material such as the preferred compound in column 3, lines 10, 30 and 65. Please note that Bojora et al. use the sulfonyl-containing compounds as an organic solvent. See Claim 1 and column 2, lines 42-53. The sulfonyl-containing compounds that can be used in Bojara et al. are compounds useful as an organic solvent. Accordingly, a person skilled in the art would only have been motivated to use the sulfonyl-containing compounds as an organic solvent.

The object of the invention described in Bojora et al. is to provide improved maximum image densities and reduced exposure and processing time, i.e. acceleration of development. This is in distinction to the object of the claimed invention which is to provide a photothermographic material that can sufficiently suppress coloration of blank portions during storage in the dark after development (see page 1, lines 3-5 of the specification). Bojora et al. are silent on the suppression of coloration of blank portions during storage. The object of the

claimed invention is achieved by using the sulfonyl-containing compound of formula (III) in combination with the phenol compound in a photothermographic material. No one skilled in the art reading Bojora et al. would have been motivated to use the sulfonyl-containing compound in order to improve the coloration of blank portions during storage. Also, no one skilled in the art would have predicted that use of the sulfonyl-containing compound improves the coloration of blank portions during storage before the claimed invention was made.

In Example 1 of Bojora et al., tetrahydrothiophene-1,1-dioxide is used as a sulfonylcontaining organic solvent. The amount is rather small. As is well known in the art, tetrahydrothiophene-1,1-dioxide is easy to volatilize. As such, most of the added dioxide will volatilize off during a heat development and therefore little dioxide will remain in the photothermographic material after the heat development. Therefore, the photothermographic material of Bojora et al. cannot suppress coloration of blank portions during storage in the dark. This is in distinction to the claimed invention, wherein the sulfonyl-containing compound of formula (III) remains and functions well in the photothermographic material after the heat development.

Also, Miura et al. are silent of the use of sulfonyl-containing compounds to improve the coloration of blank portions during storage. As such, Miura et al. do not compensate for the deficiencies of Bojora et al.

In view of the deficiencies in Bojora et al. and Miura et al., the claimed invention is thus patentable over Bojora et al. and Miura et al. and withdrawal of the rejection is respectfully requested.

In view of the above amendment, applicant believes the pending application is in condition for allowance.

Should there be any outstanding matters that need to be resolved in the present application, the Examiner is respectfully requested to contact Garth M. Dahlen, Ph.D., Esq. (Reg. Application No. 10/643,221 Amendment dated April 9, 2007

After Final Office Action of October 10, 2006

No. 43,575) at the telephone number of the undersigned below, to conduct an interview in an effort to expedite prosecution in connection with the present application.

If necessary, the Commissioner is hereby authorized in this, concurrent, and future replies, to charge payment or credit any overpayment to our Deposit Account No. 02-2448 for any additional fees required under 37 C.F.R. § 1.16 or under § 1.17; particularly, extension of time fees.

Dated: April 9, 2007 Respectfully submitted,

Marc S. Weiner

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Studies of Hydrogen-Bonded Complex Formation with p-Fluorophenol. IV. The Fluorine Nuclear Magnetic Resonance Method'

D. Gurka and R. W. Taft

Contribution from the Department of Chemistry, University of California, Irvine, California 92664. Received March 7, 1969

Abstract: Using 0.01 Mp-FC<sub>6</sub>H<sub>4</sub>OCH<sub>5</sub> as an internal reference standard in CCl<sub>4</sub>, the shielding effects of hydrogen-bonded complex formation of  $0.01\,M_P$ -FC-M-OH have been evaluated quantitatively. Using this method and reference acid, the formation constant,  $K_t$ , and limiting F nmr shifts,  $\Delta_t$  for formation of the 1:1 hydrogen-bonded complex have been obtained for 62 bases of widely different structures in CCI, at 25°. Some additional formation constants have been obtained in CCl<sub>1</sub> at -20°. In general, the yan't Hoff enthalpies of complex formation based upon  $K_t$  values at 25 and  $-20^{\circ}$  are in satisfactory accord with the calorimetric values obtained in the companion collaborative study. A linear relationship between corresponding values of  $\log K_i$  and  $\Delta$  is shown to apply with generally good precision, except for bases of large steric requirements. This empirical relationship provides evidence that the potential energy change in the formation of the hydrogen-bonded complex is approximately proportional to the standard free energy change. The F mm  $\Delta$  values are essentially independent of temperature in CCl, and are not increased in the polar solvent, ClCH<sub>2</sub>CH<sub>3</sub>Cl, indicating the formation of only hydrogen-bonded complexes (no appreciable contribution from the hydrogen-bonded ion pair, FC,H,O--+HB). The A values are interpreted as showing that a small extent of proton transfer (appearant proton transfer = 10-30%) occurs in the hydrogen-bonded complexes of p-FC<sub>1</sub>H<sub>4</sub>OH. It has been conclusively established that the aqueous  $pK_A$  and the  $\log K_t$  scales of base strengths for common functional groups are unrelated.

In this paper we report results from a collaborative affort to obtain definitive hydrogen-bond paramstars. The reference acid, p-fluorophenol, was selected for study in hydrogen-bonded complex formation with general bases in very dilute carbon tetrachloride solutions using three methods of investigation: fluorine nuclear magnetic resonance spectroscopy (F nmr), infrared spectroscopy (ir), and calorimetric determinations. Preliminary results of this collaborative work have been communicated. \*\* The F nmr method is described in detail and the results of its application to 62 bases of widely varying structure are reported herein.

Two major objectives have been explored. First, the applicability of the linear electronic energy shielding (LSEE) relationship to hydrogen-bond formation by p-FC,H.OH has been determined. Second, a detailed study has been carried out of the effects of molecular structure on base strength as measured by the formation constant of the 1:1 hydrogen-bonded complexes of p-FC,H,OH. The applicability and limitations of linear free energy and other structure-reactivity relationships to hydrogen-bonded complex formation have been determined. The results of this second major area of interest are reported in paper V of this series.

A key issue in the development of the F nmr method for the present work was the choice of a proper reference standard for evaluating the intramolecular sorcening effects of hydrogen-bonded complex formation with p-FC<sub>1</sub>H<sub>4</sub>OH. Precise formation constants,

(i) This work was supported in part by the Public Health Service, Project GM14078. We gratefully acknowledge this support and the support of the National Science Foundation which made available the nar spectrometer to the Chemistry Department,

(2) E. M. Arnett, T. S. S. R. Murty, P. von R. Schleyer, and L. Joria, J. Am. Chem. Soc., 59, 5935 (1967); paper I of series.

(3) D. Gurka, R. W. Taft, L. Joria, and P. von R. Schleyer, ibid., 89, 5937 (1967); paper II of series.

(4) R. W. Taft and L. D. McKerver, ibid., 87, 2489 (1968); 88, 4544 (1966); C. S. Glam and R. W. Taft, ibid., 89, 2397 (1967).

(5) R. W. Taft, D. Gurka, L. Joria, P. von R. Schleyer, and J. W. Rakshys, ibid., 91, 4801 (1969).

Kt, for the 1:1 complex of base and p-FC.H.OH obtained by the ir method! were used to develop a definitive F nmr reference standard. We have found that up to base concentrations of 0.02 M, the effects of base on the shielding of 0.01 M p-FC,H,OH in CCl, relative to an external reference are (within the experimental error of ±0.02 ppm) due entirely to hydrogen-bonded complex formation. However, with base concentrations above 0.02 M, the shifts based upon the external and the internal reference (cf. Experimental Section) are measurably different. The shift relative to the external reference is due to contributions from H bonding and general F nmr medium effects." Ideally, one would hope to find an internal reference having precisely the same F nmr medium effects (relative to external reference) as that of the p-fluorophenol. The F nmr shifts of p-fluorophenol measured relative to such an internal reference at base concentrations much larger than 0.02 M would then also be quantitative measures of the Hbonding shift.

The use of 0.01 M p-fluoroanisole as internal reference accomplishes this objective. Previous studies showed that the internal fluorobenzene standard nearly eliminates intermolecular F nmr medium effects and provides the intramolecular shielding effects of meta and para substituents. Since the intramolecular screening effects of the OCH, and OH substituents are very similar, p-fluoroanisole is the logical selection for an internal reference which presents to the medium a p-F atom very nearly identical with that of p-fluorophenol and its hydrogen-bonded complexes. Since 0.01 M pfluoroanisole is not expected to give (and the P nmr shielding results do not indicate any) chemical com-

(6) B. M. Arnett, L. Jorla, B. Mitchell, T. S. S. R. Murty, and P. von R. Schleyer, submitted for publication; paper III of series, (7) (a) D. F. Evant, J. Chem. Soc., 577 (1960); (b) R. W. Glick and S. Eirenson, J. Phys. Chem., 62, 1399 (1958).
(5) R. W. Taft, E. Fries, I. R. Fox, I. C. Lewis, K. K. Anderson, and O. T. Davis, J. Am. Chem. Soc., 85, 709, 1146 (1963).

binations with any of the bases employed or with the 0.01 M p-FC,H4OH, the use of this internal reference provides quantitative shifts which satisfy the hydrogenbonded complex formation equilibrium expression (cf. eq 1 following section) up to relatively high base concentrations.

Calorimetric and ir studies by our collaborators show that p-FC.H.OH is negligibly self-associated in GCl4 solutions at the concentrations employed (~0.01 M) in this study. They also find that p-fluorosnisole is a satisfactory reference for the determination of calorimetric heats of hydrogen-bonded complex formation of p-FC6H4OH. 1/8

### Experimental Section

Materials. Carbon tetraphioride and all liquid bases were distilled from calcium hydride immediately before use. Solid bases were recrystallized two to three times until agreement with literature malting points was obtained, p-Muorophenol was sublimed just before use and immediately transferred to purified carbon tetre-chloride. All bases used were obtained commercially except those given below. Samples of triphenylphosphine oxide and p-nitrophenyl methyl sulfoxide were obtained from Dr. J. R. Rekshys and Professor K. K. Andersen, respectively. N.N-Dimethyltrifiuoroacetamide was prepared by the procedure of Bissel and Finger.

Disfebutyl ether was prepared by the procedure of Brickson and

Ashton. Phenyl methyl sulfoxide was prepared by the procedure
of Leonard and Johnson. N.N.-Dimethyl-p-nitrobenzamide
(mp 95-97°; lit. 393-95°) and N.N.-dimethyl-cchloromeetamide (bp 65":(1 mm)) were prepared by the reactions of p-nitrobenzoyl chloride and a chloroscatyl chloride, respectively, with dimethyl-

amine in dry ether,

Apparatus. All spactra were obtained on a Varian HA-60
spectrometer equipped with a Varian 1024 computer of average
translans (CAT) and NMR Specialties heteronuclear decoupler,

transants (CAI) and NMM Specialites deteronucies decoupler, Model SD50-B. The temperature was maintained at either -20 or 25° using the temperature-control probs.

All measurements were made with solutions of purified carbon starachloride containing 0:01 M p-FC<sub>1</sub>H<sub>1</sub>OH and 0:01 M p-FC<sub>1</sub>H<sub>1</sub>OH<sub>2</sub>OH<sub>3</sub> (the internal reference). The nurr tubes contained an inserted capillary of 28 wt % 1,1,2;2-tetrafluoro-3,3,4,4-tetrachloro-syclobulane in CCI, as external reference, 1. The ring protons were decoupled and about five CAT some were employed for each decoupled and about five CAT scans were employed for each "fitration" point. The CAT recorder was calibrated to agree with direct read-out within ±1 cps. All spectra were run at a sweep width of 250 cps and a sweep time of 100 acc.

For base concentrations up to ca. 0.02 M, the same F nmr shifts, a = 0.02 ppm), is obtained with either the internal or external references. From 0.02 to 0.7 M base only the internal reference gives satisfactory results (cf. following section).

### Data Treatment

"Equilibrium Expression. For the formation of a 1:1 hydrogen-bonded complex between p-FC,H,OH and general base, B, the following equilibrium expression applies.

p-PC,H,OH + B → p-FC,H,OH--B

The formation constant,  $K_{t_1}$  is shown in eq 1 where  $K_t = (\delta/\Delta)A_0/[A_0[1 - (\delta/\Delta)]][B_0 - (\delta/\Delta)A_0] \quad (1)$ 

 $A_0 = \text{initial (total) } p\text{-PC}_1H_1OH concentration, <math>H_0 =$ initial (total) base concentration, and  $\delta =$ the observed time-average F nmr shift in parts per million (ppm) for the equilibrium mixture relative to that for 0.01 M p-FC,H4OH in CCl4. Unless otherwise specified the reference standard is internal 0.01 M p-FC.H4OCH1.

(9). R. B. Bluel and M. Finger, J. Org. Chem., 24, 1236 (1939). (10) J. L. E. Erickson and W. H. Abhton, J. Am. Chem. Soc., 63, 1769 (1941).

(11) N. J. Leonard and C. R. Johnson, ibid., 54, 3701 (1962). (12): A. V. Kirsanov, Zh. Obshell. Khim., 23, 223 (1953); Chem. Abstr., 48, 2635a (1954).

A - the limiting F nmr shift (ppm), i.e., the shift of the completely formed complex relative to uncomplexed p-FC,H,OH,

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Validity of Equilibrium Expression. When the value of K is known from an independent determination, e.g., the ir method, eq 1 may be rigorously confirmed by the demonstration that the values of A calculated for each 8 value are a precise constant, i.e., independent of  $B_0$ . Alternatively, the observed and caloulated values of a at each base concentration over the range of 30-80% complex formation may be shown to be in agreement (to within the combined experimental errors), based upon the use of the best value of A. Using the K, values provided by the collaborative work of Arnett, Murty, Schleyer, and Joris, this condition was found to hold in every instance. Typical results are illustrated in Table I.

Table I. Typical Fits of F Nmr "Titration Curve" to Equilibrium Eq 1 at 25° in CCl.

			<u> </u>
10°A, M	10³ <i>B</i> <sub>0</sub> , <i>M</i>	Obsd `	f, ppmCalcd
	N,N-Dimethyl	acetamide*	* -
1.126 -	0,409	0.71	0.70
1.052	0.782	1.22	1.25
0,126	0.816	1,25	1,28
1.052	1.174	1,62	1,62
1.126	1 223	1.65	1.66
1 .052	1.565	1.94	1.93
1.126	1.63	1.96	1.93
7.126	2:04	2.14	2.13
', <b>'</b>	Ethyl Ace	tates	
0.934	2,52	0.40	0,40
0.934	4.20	0.59	0.59
0.934	8,40	0.90	0:90
0,934	12,59	1.08	1,09
0,914	15.8	1.22	1,22
0.934	.21.0	1,33	1.31

 $^{5}K_{1} = 242 \pm 6$ ;  $\Delta = 2.86$  ppm.  $^{5}K_{1} = 12.0 \pm 0.2$ ;  $\Delta = 1.85$ 1. 1.1.11.65

In view of its widely demonstrated applicability, eq 1 may also be accepted as valid for those bases for which an independent K, value is not available. The syaluation of the K, and A value for these bases from a values observed over a range of 30-80% complex formation is described subsequently."

Validity of the Internal Reference. Identical shifts were observed up to 0:02 M concentrations of all bases with either the external reference or internal reference. Consequently, the demonstration of the validity of eq 1 with strong bases at concentrations less than 0.02 M shows rigorously that the shifts relative to external reference are due entirely to hydrogen-bonded complex. formation (no F amr medium effects). The data for N,-N-dimethylacetamide in Table I are typical. Table II lists additional results for strong bases for which eq 1 holds at base concentrations less than 0.02 M. The precision of eq 1 is demonstrated by the precision of the A value obtained (Table II gives the mean value of Δ(ext.zu) from Δ values calculated at each base concentration for the indicated range; deviations from the mean value are entirely random, showing no trends

Table'II also lists corresponding values of  $\Delta^{(int rel)}$  obtained from direct measurement using 0.01 M internal

Table II. Agreement between Calculated A Values from Measurements with Partial Conversion Using the External Reference and Observed A Values at 97-99 % Conversion Using 0.01 M Internal p-FC,H,OCH;

Beso	10'B <sub>0</sub> , M	K <sub>i</sub> *\$ " (ir)	Variet(ex) tel)	∆obed(lat rel)
Dimethyl sulfoxido 2,6-Dimethyl-y-pyrone N,N-Dimethylacetamide N,N-Dimethylformamide Triethylamine Pyridine	0.48-1.94	346 ± 8	2.70 ± 0.01	2.74 ± 0.02
	0.30-1.51	318 ± 18	3.07 ± 0.05	2.99 ± 0.02
	0.41-1.63	260 ± 12	2.83 ± 0.03	2.86 ± 0.02
	0.67-2.05	116 ± 3	2.71 ± 0.01	2.72 ± 0.02
	0.64-1.92	85.2 ± 1.9	2.67 ± 0.02	2.66 ± 0.02
	0.69-1.93	76.2 ± 1.1	2.57 ± 0.03	2.51 ± 0.02

Table III. Comparison of Independently Determined Formation Constants, Ki, for p-FC:H;OH-Base Complexes, CCi., 25.0°

Base	Ir method	F nmr method	Calorimetria method	
Dimathyl sulfoxide	346 sts 8	338 🖶 7	1	
N,N-Dimethylacetamide	260 ± 12	242 ± 6		
4-Mathoxypyridine	131 ± 4	139 土 2		
N.N-Dimethylformamide	115 ± 3	115 业 2	122 ± 9	
4-Methylpytldine	109 ± 5	107 # 2	102 ± 7	
Triethylamine	$85.2 \pm 1.9$	82 ± 2		
Pyridine	$76.2 \pm 1.1$	76 ± 1	74 🕁 5	
Quinoline	72.3 = 1.0	$71 \pm 3$	73 ± 5	
Cyclohaxanone	$20.5 \pm 0.7$	$21.4 \pm 0.5$		
Tetrahydrofuran	17.7 ± 0.5	18.4 ± 0.5	$19.4 \pm 1.0$	
2-Butanons	15,6 ± 0,5	15.1 =4 0.3	17.0 ± 1.0	
Ethyl acetato	$12.3 \pm 0.3$	12.0 ± 0.2	$13.0 \pm 0.7$	
Diethyl ether		$9.5 \pm 0.4$	10.3 🖶 0.3	

p-FC.H.OCH, as reference and high base concentrations (0.1-0.7 M) such that the hydrogen-bonded complex is 97-99% formed (with correction to 100%). The substantial agreement shown in Table II between corresponding values of A(lat rat) and A(ext rat) for bases of a wide variety of types establishes the validity of the internal reference procedure. The result is made rigorous by the finding in the collaborative ir determinations that values of K: are constant (within their precision measures) for similar ranges of base concentrations. Consequently, the agreement between  $\Lambda^{(axt rai)}$  and  $\Lambda^{(int rai)}$  cannot be an accident of coincidental cancellations of F nmr shielding and K; medium effects.

Further confirmation comes from the fact that eq 1 is also valid for those weak bases which require concentrations appreciably greater than 0.02 M in order to zohieve 30-80% complex formation. Using shifts based upon 0.01 M internal p-FC.H.OCH, and the independently determined K, values from the ir method, the results for weak bases are found to follow eq 1 to within the combined experimental errors of observed and calculated 8 values. The results given in Table I for ethyl acetate are typical in this regard.

Final confirmation is shown by the excellent agreement obtained between the independently determined K, values, i.e., values based upon only F nmr, ir, or calorlmetric determinations. The method employed for independent F nms determination of the  $R_t$  value is given in the following section. The ir and calorimetric methods are reported in the companion paper.8 A summary showing the excellent agreement obtained by the three methods is given in Table III.

We conclude that 0.01 M p-PC<sub>4</sub>H<sub>4</sub>OCH<sub>2</sub> as an internal reference in CCL, may be used generally with most types of bases (up to concentrations of  $\sim 0.7 M$ ) to obtain the F nmr shielding effects for hydrogen bonding with 0.01 Mp-FC,H,OH,

Independent F Nmr Method for Determination of  $K_t$ . The shifts, &, obtained for the range of given base concentrations required to achieve approximately 30-80% complex formation, were analyzed by computer16 in terms of eq 1 to give best value of the formation constant, Ki, and the corresponding limiting shift, A. The analysis is subject to the following considerations in light of the results presented above. The accumulation of error in one parameter necessarily leads to error in the other parameter (i.e., in both  $K_i$  and  $\Delta$ ), and, in general, such errors are not readily apparent in a poorer precision of over-all fit of the 8 values to eq 1.3 However, this difficulty can be largely overcome by the direct measurement of  $\Delta$  in independent experiments. Computer analysis of the  $\delta$  values with the fixed value of  $\Delta$ gives a best value of K, which is relatively precise (cf. Table III) and the fit of data is essentially to the experimental error of individual points (±0.02 ppm). All of the bases employed in this study for which apparently satisfactory best values of  $\Delta$  were obtained (cf. following section) gave such results.

Evaluation of Best Values of  $\Delta$ . Three different determinations of A were developed from which a best value was selected. First, values referred to as  $\Delta^{titr}$ were obtained from the 8 values for the range 30-80% complexing (relatively low base concentrations) using computer analysis of best fit to eq 1. If an independently determined K, value was available (i.e., from it determination) this value was used as a constraint in the computer analysis for results summarized in Table V. Otherwise, the Attir value is that obtained when the computer is free to select best values of both A and K, (as employed to obtain the completely independent F nmr results given in Table III, or as employed in the absence of an ir determination of  $K_i$ ).

Second, values of A were obtained at 25° by direct measurement at the high base concentration (0,1-0.7 M)which gives 97-99% complex formation (corrected to 100%). Third, a procedure similar to the second was

(13) R. W. Taft, G. B. Klingensmith, and S. Ehrenson, J. Am. Chem. Soc., 67, 3620 (1965).

Table IV. Evaluation	on of Annie by Three Method	4		K
Hase no.	∆altr ∆altr	" Ditt tef)	· Δilos seti	. Abert
1		3.71 ± 0.02	•	$3.71 \pm 0.02$
1 2	$3.58 \pm 0.03$			$3.58 \pm 0.03$
3	$3.40 \pm 0.09$	$3,40 \pm 0.02$	3,34 ± 0.02	$3.37 \pm 0.05$
4	3,00 ± 0.01	3.00 ± 0.01		3.00 ± 0.02
<b>7</b> 5	$2.90 \pm 0.04$	2.97 📥 0.02	$3.00 \pm 0.02$	$2.96 \pm 0.04$
i: 6	$3.07 \pm 0.04$	$2.98 \pm 0.02$	$2.89 \pm 0.02$	$2.98 \pm 0.05$
6 7 8 9	2.83 仕 0.02	$2.91 \pm 0.01$	2.84 ± 0.02	2.86 由 0.04
i . 8	2.82 ± 0.01 2.77 ± 0.01 2.71 ± 0.01	2.74°± 0.02	$2.73 \pm 0.02$	. 2.77 ± 0:03
9	$2.77 \pm 0.01$	$2.75 \pm 0.02$	2.66 ± 0.02	2.74 ± 0.02 2.72 ± 9.02
	$2.71 \Rightarrow 0.01$	2.72 ± 0.02 2.74 ± 0.02	2.66 ± 0.02	2.71 並 0.03
11	· 2.71 == 0.01	2.74 == 0.02	2.00 50 0.02	2.71 ± 0.03 2.71 ± 0.04
11 11 12 12	$2.68 \pm 0.03$	2,75 ± 0.02		2.70 ± 0.02
ii.	$2.69 \pm 0.02$	$2.71 \pm 0.02$ $2.69 \pm 0.02$	•	2.70 ± 0.02
14	$2.71 \pm 0.01$	2.09 22 0.02		2.67 ± 0.02
15 15 16	$2.67 \pm 0.01$	$2.67 \pm 0.02$	0.70 0.00	$2.66 \pm 0.02$
16	2.64 st 0.02	2.65 ± 0.02	$\begin{array}{c} 2.70 \pm 0.02 \\ 2.52 \pm 0.02 \end{array}$	2.00 ± 0.02
17	$2.59 \pm 0.02$	$2.60 \pm 0.02$	2.32 = 0.02	$2.54 \pm 0.04$ $2.52 \pm 0.02$
18	$2.53 \pm 0.01$	$2.51 \pm 0.02$		2.50 ± 0.06
18 19 20	$2.50 \pm 0.06$	a at 1 a a ma	ŕ	
20 عراج	$2.46 \pm 0.01$	2.91 ± 0.02		2.49 ± 0.03 2.45 ⇒ 0.02
7.1	2.44 📥 0.01	$3.45 \pm 0.02$		2.41 ± 0.04
22		2,41 ± 0.02	n an al. Dan *	
23 24	$2.40 \pm 0.03$	2.34 注 0.02	2.40 ± 0.02 ° 2.39 ± 0.02	$2.40 \pm 0.03$ $2.40 \pm 0.03$
24	$2.41 \pm 0.04$	$2.34 \pm 0.02$	40.0 m vc.4	2.33 ± 0.05
L 25	2.30 ± 0.03	$2.37 \pm 0.02$		2.32 ± 0.03 '
26	2.32 # 0.05			2.29 ± 0.02
(a. <b>41</b>	2.29 ± 0.02 2.21 ± 0.03		2.23 ± 0.02	2.23 ± 0.04
28 29	2.21 ± 0.03 2.22 ± 0.04	.•	#1#D == 0.0#	2.22 = 0.04
30	2.22 = 0.04		$2.16 \pm 0.05$	2,16 = 0.05
31	2,18 🕁 0.04	$2.11 \pm 0.02$	2.18 ± 0.02	2.16 ± 0.03
32	$2.12 \pm 0.01$	2.17 4 0.02	#120 -1 010E	2.15 ± 0.03
33	$2.14 \pm 0.02$	1	$2.10 \pm 0.02$	2.12 ± 0.02
34	ማ 10 👊 ስ በዓ			$2.10 \pm 0.03$
ic 35	2.07 ± 0.03 2.00 ± 0.00			$2.07 \pm 0.03$
9. 36	$2.00 \pm 0.00$		2,03 = 0.02	$2.02 \pm 0.02$
36 37	$2.00 \pm 0.05$		2.02 ± 0.02	2.01 at 0.03
38	$2.02 \pm 0.00$		$1.97 \pm 0.02$	2.00 # 0.02
69° 39	1.91 st 0.04		$2.07 \pm 0.02$	1.99 ± 0.05
∫6· 40	1.92 ± 0.03 1.88 ± 0.03		$1.92 \pm 0.02$	$1.92 \pm 0.02$
41	1.88 = 0.03	•	1.91 ± 0.02	1.90 ± 0.02
42			I. N.K ±1 0.02	$1.88 \pm 0.02$
Gr. 43	$1.85 \pm 0.03$		1,90 ± 0.02 1,87 ± 0.03 1.86 ± 0.02	1.88 === 0.03
44	1.84 土 0.04	1.89 ± 0.02	1.87 de 0.03	1.88 ± 0.02
49	1.83 ± 0.01		1.86 土 0.02	1.85 ↔ 0.02
46	$1.84 \pm 0.02$	•	` '	1.84 # 0.02
47	1.78 🖶 D.OS	•	1,87 📥 0.02	1.83 📥 0.05
(J. 48	2.17 土 0.2		1.75 ± 0.02	•
49	2.17 ± 0.2 1.69 ± 0.06	•	$1.72 \pm 0.02$	1.71 == 0.03
50	1,64 ± 0.04		1.73 4 0.02	$1.70 \pm 0.05$
4: 31	1.70 ab 0.04			1.70 da 0.04
52	$1,60 \pm 0.02$		$1.63 \pm 0.02$	$1.62 \pm 0.03$
· 53 ·	$1.48 \pm 0.4$		$1.42 \pm 0.02$	1.45 == 0.04
S4	$1.40 \pm 0.05$		1.45 == 0.02	1,45 由 0.04
55	$1.12 \pm 0.10$		$1.00 \pm 0.10$	$1.10 \pm 0.10$
1.7.7		~~~~		

In parts per million. Base number refers to the designation given in Table V.

employed at a temperature of -20° where the equilib-Hum is shifted in favor of further complex formation.
The third procedure was especially useful for bases too

weak to utilize the second procedure at 25°. Table IV illustrates the generally good agreement ob-Hillined by these methods. Since the potential sources of Pror for Atter are quite different then those for A (lat rai). the general agreement achieved offers strong additional upport for the conclusions of earlier sections. Except f 2,6-dimethyl-y-pyrone, trimethyl phosphate, and lifficity sulfoxide (bases numbered 6, 9, and 11) there potent to be no significant differences between  $\Delta^{(lateral)}$  and  $\Delta^{(lateral)}_{-20}$ . Since  $\Delta^{(lateral)}$  is subjected to fror if the base is appreciably self-associated at the elatively high concentrations (several tenths molar) employed in this method of determination, it is probably significant that  $\Delta_{20}^{(i_1,i_2)}$  is smaller than  $\Delta_{22}^{(i_1,i_2)}$  for each of these three bases. These three bases are of the polar type which tend to show the greatest self-association, and such self-Masociation should be greater at -20 than 25°. Competition with hydrogen bonding would thereby reduce the apparent value of A. Values of  $\Delta_{best}$  for these three bases were therefore taken from the results at 25° only. For all other bases, values of Abest were selected either by averaging, or, if precision measures were substantially different, by weighting in favor of the more precise individual values.

It is also significant that self-association of a base would be expected to cause deviations from the equilibrium expression, eq 1. Consequently, the fits of the Table V. Summary of Formation Constants and Limiting Shifts for Hydrogen-Bonded Complex Formation with a FC.H.OH, CCl., 25°

No.	Base	Δ, ppto	Kı, M-1 4	Log K <sub>f</sub>
·	Hexamethylphosphoramide	3.71 ± 0.03	3600	3.56 ± 0.06
1	Triphenylphosphine oxide	3.58 ± 0.03	1456 ± 80	$3.16 \pm 0.06$
2		9 99 I N NE	650 ± 90	2.81 ± 0.06
3	4-Dimethylaminopyridine	$3.00 \pm 0.02$	261 ± 5	$2.42 \pm 0.01$
4 5 6	Tetramethylurea	2 06 4 0.04	167 🚣 16	$2.22 \pm 0.04$
5	N,N-Dimethylbenzamide	2.98 = 0.05	318 ± 18	2,50 土 0.03
б	2,6-Dimethyl-y-pyrone	7.50 m 0.03	242 ± 6	$2.38 \pm 0.01$ $2.14 \pm 0.01$
7 8	N,N-Dimethylacetamide	2.00 == 0.04	139 ± 2	$2.14 \pm 0.01$
8	4-Methoxypyridine	2.77 ± 0.03	250 ± 8	$2.40 \pm 0.01$
ğ	Trimethyl phosphate	2.74 == 0.02 6.70 J D D2	115 ± 2	$2.05 \pm 0.01$
10	N,N-Dimethylformamide	3.00 ± 0.02 2.96 ± 0.04 2.98 ± 0.05 2.86 ± 0.04 2.77 ± 0.03 2.74 ± 0.02 2.72 ± 0.02 2.71 ± 0.03	338 並 7	$2.53 \pm 0.01$
11	Dimethyl sulfoxide	$2.71 \pm 0.03$	Strate G	2.07 ± 0.01
12	N,N-Dimethyloyolohexylamine	$2.71 \pm 0.04$	118 ± 2	2.03 ± 0.01
13	4-Mathylpyridina	$2.70 \pm 0.02$	107 土 7	2.03 ± 0.01
14	Dinhenyl sulfoxida	$2.70 \pm 0.02$	$106 \pm 2$	\$,03 ± 0.01
14 15	Phenyl methyl sulfoxido	2.67 to 0.02	141 世 4	$2.15 \pm 0.01$
16	Triethviemins	$2.66 \pm 0.02$	85 ± 2	1.93 ± 0.01
17	N,N-Dimethyloxida N,N-Dimethyloxida N,N-Dimethyloxida A-Methylpyridina Diphenyl sulfoxida Phenyl methyl sulfoxida Triethylamina N,N-Dimethyl-n-propylamina Z-n-Butylpyridina Tri-n-butylamina Pyridina Flavona N,N-Dimethyl-hloroscetamida Quinolina Benzylamina	2,54 ± 0.04	95 共 1	1.98 ± 0.01
10	7. m. Wittelmoridina	2,52 de 0.02	$76 \pm 2$	1.88 ± 0.01
18	Marin Marianinas	$2.50 \pm 0.05$	37 ± 3	$1.57 \pm 0.03$
19	I [-i-but inte-	2.49 ± 0.03	76 🛨 1	1,88 由 0.01
20 21	Pyriamo	$2.45 \pm 0.02$	98 ± 6 48 ± 5 71 ± 3	$1.99 \pm 0.03$
21	Figure Flavors N.N-Dimethylchioroscetamids Quinolins Benzylamine N.N-Dimethylbenzylamine p-Nitrophenyl methyl sulfoxides Cyclopropylamines N.N-Dimethyl p-nitrobenzamide Trisilylamines p-Dimethylaminobenzonitriles Fropargylamine Acetylferrocens Cyclohexanene p-Methokyavetophenones 2-Bromopyridines 2-Butanone Di-t-butyl ether Teirelydrofuran	$2.41 \pm 0.04$	48 ± 5	$1.68 \pm 0.04$
22 23 24 25 25	M'M-Diffictulateurotescersumos	2 40 - 0 03	71 🖦 3	1.85 == 0.02
23	Qumotini	2 40 -4 11 113	56 ± 2	1.75 由 0.01
24	Benzylamine	2 33 -6 0 04	38 ± 3 38 ± 5 44 ± 2 48 ± 6	1.58 ± 0.03 1.58 ± 0.05
25	N.N. Dimemyloentylumine	2 32 + 0.05	38 🕁 5	1.58 📥 0.05
25	b-Mittobustar merual annexton.	2 20 0 02	44 ± 2	1.64 土 0.02
27	Cyclopropylamine	70.0 T EC 0	48 6 6	$1.62 \pm 0.03$
28	N.N.Dimethyl p-nitrobenzamice	2.23 20 0.04	17 ± 1 24 ± 3	$\begin{array}{c} 1.62 \pm 0.05 \\ 1.23 \pm 0.03 \end{array}$
29 30	Trinityiamina.	2 15 ± 0 04	24 + 3	$1.38 \pm 0.05$
30	p-Dimethylaminobanzonictile.	2,10 == 0.03 4 16 ± 0.01	30 ± 1	1.48 ± 0.02
31	Propargylamine	2.10 = 0.03	4 ± 1	1.64 土 0.01
32 33	Acetylierroccus	2,13 at 0.03	$21.4 \pm 0.5$	1,33 ± 0.01
33	Cyclohexanona -	2,12 = 0.02	25 4 1	1.40 ± 0.02
34 35 36 37 38	p-Methoxyacetophenone	X.10 == 0.03	8.8 # 0.5	0.94 ats 0.03
35	2-Bromopyridina	2.0/ 23 0.03	15.1 = 0.3	1.18 ± 0.01
36	2-Butanons	2.02 = 0.02	5.1 ± 0.4	0.71 🕳 0.03
37	DI-A-butyl ether	2.01 # 0.03	18.4 ± 0.5	1,26 ± 0,02
38	Teirshydrofuren	2.00 = 0.02	40 1 1 1 7	$1.26 \pm 0.03$
39	3-Bromopyridine	1.99 # 0.03	18.4 ± 0.5 18.1 ± 1.2	1.13 ± 0.01
40	Acatophanona	$1.92 \pm 0.02$	13.0 == 0.3	1:10 ± 0.01
40 41	p-Methoxybenzaldehyde	2.10 ± 0.03 2.07 ± 0.03 2.02 ± 0.03 2.01 ± 0.03 2.00 ± 0.02 1.99 ± 0.02 1.92 ± 0.02 1.88 ± 0.03 1.88 ± 0.03 1.88 ± 0.03	$12.6 \pm 0.2$	1,05 土 0,02
42	Acatonitrile*	$1.88 \pm 0.02$	$11.2 \pm 0.6$	0.98 占 0.01
43 44 45	Diothyl other	1,88 曲 0,03	9.5 ± 0.2	
44	β-Cyanosthyl ethyl other	1,88 ± 0.03	$11.2\pm0.5$	1.05 ± 0.07
43	Ethyl acetate	$1.85 \pm 0.02$	$12.0 \pm 0.2$	$1.08 \pm 0.01$
46	Pyrimidino*	$1.84 \pm 0.02$	$22.5 \pm 0.5$	$1.05 \pm 0.02$
47	p-Methoxybenzonitrile	1.84 ± 0.02 1.83 ± 0.05 (1.75) <sup>6</sup>	9.8 # 0.4	0.99 ± 0.02
48	N,N-Dimethylaniline	(1.75)*	2.7 ± 0.2	0,43 ± 0.03
49	Banzonitrile	$1.71 \pm 0.03$	$6.1 \pm 0.6$	$0.79 \pm 0.04$
50	Benzaldehyda	1.70 ± 0.05	$6.8 \pm 0.2$	$0.83 \pm 0.01$
	Dibenzyi ether	1.70 由 0.04	5.3 ± 0.2	$0.72 \pm 0.02$
51		$1.62 \pm 0.03$	$5.6 \pm 0.2$	$0.75 \pm 0.0$
52	3,5-Dichloropyridina	1.45 ± 0.04	$3.9 \pm 0.1$	$0.59 \pm 0.01$
53	$\beta_i\beta_i\beta$ -Triffucroethylamine	1.45 ± 0.04	$10.2 \pm 0.6$	$0.71 \pm 0.07$
54	Dioxano	1.10 ± 0.10	$1.3 \pm 0.2$	$0.11 \pm 0.0$

<sup>\*</sup>With statistical correction. \ Cf. comments in Experimental Section. \ Values of K; are based only upon a single determination of the A value; of, Table IV. \*At 25°.

present data to eq 1 provide no evidence of appreciable self-association in CCl<sub>4</sub> of the bases of Tables IV and V at the concentration levels employed to give 30-80% complex formation with 0.01 Mp-FC<sub>4</sub>H<sub>4</sub>OH<sub>4</sub>.

The results for N,N-dimethylaniline (cf. Table IV) are exceptional, as were those obtained with n-butylamine. For the latter, eq I is well fitted by the best parameters:  $\Delta_{28}^{\text{dis}} = 2.56 \pm 0.03$ ,  $K_l = 129 \pm 6$ , However, values of  $\Delta_{23}^{\text{fin}} = 2.74 \pm 0.02$  and  $\Delta_{-80}^{\text{fin}} = 3.01 \pm 0.02$  were obtained. These results remain anomalous, although there is evidence that N,N-dimethylaniline has two acceptor sites, 14 which

(14) L. L. Perstandig, Tetrahedron Letters, 1235 (1963).

might explain its behavior. In the case of n-butyl-amine, alternate possibilities include higher order complexing at high base concentrations (involving N-H proton donation), or absorption of atmospheric carbon dioxide (which was not rigorously excluded). Our results with amines in general need to be confirmed by ir studies. We do not believe that our results are seriously affected by reaction between strongly basic amines and CCl4 for the following reasons: (a) no precipitates were observed during the time required for the F nmr measurements (although these do form on longer standing), (b) the F nmr shifts are reproducible for up to 5 hr after preparation of the solutions and these shifts obey eq 1.

Table VI. Formation Constants at -20° and Enthalpies for Hydrogen-Bonded Complex Formation with p-FC.HiOH.

MNO.	KC***	K(110	-AH(°)	-Δ <u>H</u> ι° •	-∆Hi°s
3-Bromopytidina 3-Bromopytidin	107 立 7 103 ± 8 64 ± 6 49 ± 2 40 ± 3 22.0 ± 0.5 ine 9.8 ± 0.5	18.1 18.4 15.1 12.0 10.2 5.3 2.7	5.8 5.7 4.8 4.7 4.6 4.7 4.2	6.2 ± 0.2 5.6 ± 0.1 4.0 ± 0.4	5,8 ± 0,1 5,2 ± 0,1 4,7 ± 0,1 5,1 ± 0,1 4,6 ± 0,1 4,0 ± 0,1

In CCli; measured in kilouslories/mole. From van't Hoff equation for K, at -20 and 25°; estimated uncertainty, ±0.3 kcal-Colorimetric value at 25°; cf. ref 6. Obtained by pure base method; cf. ref 6.

# Results

Table V summarizes the best values of the formation Constants, K<sub>1</sub>, for hydrogen-bonding complex formation of p-FC<sub>1</sub>H<sub>2</sub>OH with 55 different bases in CCl<sub>4</sub> solution at 25°. Also given are corresponding best values of Δ, the limiting F nmr shift in parts per million (ppm) between complexed and uncomplexed p-FC<sub>1</sub>H<sub>4</sub>OH.

Ji-Table VI summarizes the best values of K<sub>1</sub> obtained with eight bases in CCl<sub>4</sub> at -20°, together with values

with eight bases in CCl<sub>4</sub> at  $-20^\circ$ , together with values of the van't Hoff enthalpy of complex formation (based upon  $K_1$  values at -20 and  $25^\circ$ ),  $-\Delta H^\circ$ <sub>1</sub>, and corresponding calorimetric values of  $-\Delta H^\circ$ <sub>1</sub> (25°) obtained in the collaborative study. Values of  $K_1$  at  $-20^\circ$  are based upon concentrations corrected for the density thange of CCl<sub>4</sub> between 25 and  $-20^\circ$ .

#### Discussion

The molecular composition of the hydrogen-bonded domplexes of p-FC<sub>4</sub>H<sub>4</sub>OH under the conditions of the present investigation is shown rigorously to be 1:1. The fits of 8 values to the equilibrium expression 1 are generally within the experimental errors of the individual measurements for a critical region of 30 to 80% bomplex formation. Similar fits of the ir and calcifiatric data to corresponding equilibrium expressions for 1:1 complex formation have also been found, continuation, AH<sub>2</sub>, has nearly the same value in CCl<sub>1</sub> at high dilution as in the pure base as solvent. continuation that the 1:1 molectular composition does not change with dramatic interesses in the base to acid ratio.

Comparison of the values of the formation constant, A, determined independently by the three methods (Table III) shows that there are no serious deviations, agreement frequently being within the combined uncertainty limits. In view of the appreciable variety of base types for which this result has been obtained, the F mmr method developed herein may be accepted as a generally useful and precise method for the determination of formation constants.

The results given in Table VI provide a further indication of the high degree of internal consistency in the results obtained by the three methods of investigation employed in our collaborative efforts. There is generally satisfactory agreement between the values of the chihalples of complex formation,  $-\Delta H_I^{a}$ , obtained by the van't Hoff squation from  $K_I$  values at -20 and  $25^{\circ}$ , and the direct calorimetric measurements at  $25^{\circ}$ . Calformetric measurements of  $-\Delta H_I^{a}$  over a smaller temgerature interval (12, 25, and  $34^{\circ}$ ) show no measurable temperature dependence (i.e.,  $\Delta C_P$  for complex formafield is indicated to be small). The formation constants,  $K_l$ , summarized in Table V are not of uniform quality. Values of  $K_l$  which are based upon a single determination of the  $\Delta$  value (cf. Table IV) and which have not been confirmed by an independent method should be regarded as having potential uncertainty beyond their indicated precision measures (cf. footnote c of Table V).

Regarding the Linear SEE Relationship. This relationship involves a linear correlation of the F nmr shift between product and reactant states for a series of closely related chemical reactions with the corresponding thermodynamic standard free energy or enthalpy change. It has been observed in several chemical reactions quite different in character from the present one, in particular, for reactions involving strong rather than weak bond formation. In the systems studied previously the relationship has been found to hold over significant ranges of shifts and thermodynamic properties, and the latter have been taken to provide a reasonably quantitative estimation of the potential or electronic energy change for the reactions.

Figure 1 plots the log K<sub>l</sub> vs. Δ values for 48 bases of Table V. The number given in Figure 1 refers to the base as designated in Table V. An approximate linear relationship of useful precision (the standard deviation of log K values is 0.09 with a root mean square of 1.76) is obeyed. The correlation covers oxygen, nitrogen, and sulfur bases of a wide variety of types with apparently equal precision. Seven bases from Table V show deviations greater than three times the SD of Figure 1 and have been excluded: (5) N,N-dimethylbenzamide, (11) dimethyl sulfoxide, (19) tri-n-butylamine, (29) tri-allylamine, (35) 2-bromopyridine, (37) di-t-butyl ether, and (48) N,N-dimethylamiline. The anomalous behavior of the last amine is discussed in the Experimental Section.

Bases 19, 29, 35, and 37 quite plainly have the common structural feature of very high steric requirements. It is also probably significant that the additional bases of large steric requirements show smaller deviations in the same direction, i.e., triethylamine, 16; diethyl other, 43; dibenzyl ether, 51. The linear correlation of Figure 1 must be limited, therefore, to bases of small or moderate steric requirements. Deviations for the bases of large steric requirements are all in the direction indicating the P nmr shift has a much smaller dependence upon steric requirements than does the standard free energy change of complex formation. A similar result has been observed by Dr.A. A. Grey in the formation of Lewis acid-base adducts with p-fluorophenylboron dichloride. If The significance of these findings is currently under further study.

(15) A. A. Grey, Ph.D. Thesis, University of California, Irvine, June 1967.

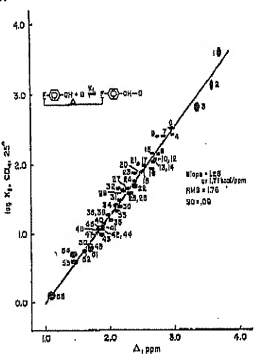


Figure 1. Linear SEE relationship for hydrogen-bonded complex formation with p-FC<sub>6</sub>H<sub>4</sub>OH, CCl<sub>4</sub>, 25°. Numbers refer to bases as designated in Table V.

The deviations of bases 5 and 11 are anomalous and no satisfactory explanation can be given presently. In the case of dimethyl sulfoxide the smaller than expected  $\Delta$  value might be interpreted as reflecting some O-S interaction in the hydrogen-bonded complex, i.e.

However, this possibility is not supported by the behavior of both phenyl methyl sulfoxide and diphenyl sulfoxide which are essentially "normal" in Figure 1.

In Table VII are presented some additional best

Table VII. Values of  $\Delta$  and Estimated Log  $K_1^{10}$ ° for Some Additional Bases

No.	Basa	Δ, ppm	Log K <sub>1</sub> 15' (est)
56	Cl=(n-CrH11)1N+	4,304	~4.0
57	Tetramethylguanidine	3.7B	3,48
58	N-Methylpyrrolidons	2.91	2.37
59	Tetramethylene sulfoxide	2.80	2,22
60	p-Dimethylaminobenzaldehyde	2,30	1.56
61	N,N-Dimethyltrifluoroacetamide	1.97	1.19

<sup>·</sup> Unpublished result of Dr. Jon Lipowits.

values of  $\Delta$  obtained by either the second or third procedure (cf. Experimental Section). Using the correlation of Figure 1, estimated values of  $\log K_1^{\rm mio}$  for these bases have been obtained and are also listed in Table VII.

The generally successful correlation of Figure 1 has interesting and significant implications with respect to

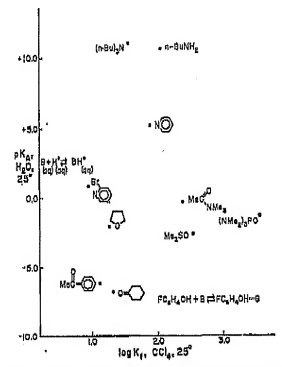


Figure 2. Comparison of base strengths, aqueous H+ vs. p-FCe-HoH.

the standard free energy change of hydrogen-bonded complex formation, AGIo. First, by the basic hypothesis of the SEE relationship, the correlation of Figure I implies that  $\Delta G_t$  is approximately proportional to the potential energy change of formation of the hydrogenbonded complex. It is particularly significant that the correlation includes bases with substantial variations in the entropies of complex formation. Thus, it is suggested that the kinetic energy terms (-RT in  $\pi Q$ ) included in  $\Delta G_i^{\circ}$  are of the special kind which, at least approximately, tend to parallel corresponding potential enorgy changes. 16 It is to be noted that the proportionality constant between the standard free and potential energies is implied to be temperature dependent by the slopes of log  $K_t$  vs.  $\Delta$  (1.71 keal/ppm at 25° and 1.92 keal/ppm at  $-20^\circ$ ). Secondly, since the F nmr shift,  $\Delta$ , has been postulated to be a measure of the "apparent per cent proton transfer" (≥100 \( \Delta/14 \) in the hydrogenbonded complex, the correlation of Figure 1 indicates that  $\Delta G_i$ ° must also be an approximate "measure" of the extent of proton transfer in the complex.

The present F nmr study has provided additional new evidence defining the character of the hydrogen-bonded complexes of  $p\text{-}FC_1H_4OH$ . It is indeed conceivable in terms of previous concepts and discussions of hydrogen bonding "that the observed values of  $\Delta$  may pertain to a complex mixture of very rapidly interconverting states having appreciably different energy contents and extents of proton transfer (e.g., the hydrogen-bonded complex, the hydrogen-bonded ion pair, and perhaps other states). We believe this possibility is excluded by the present finding that the F nmr shift,  $\Delta$ , is the same at -20 as

(16) R. W. Teft in M. S. Newman "Steric Effects in Organic Chemistry," John Wiley & Sons, Inc., New York, N. Y., 1935, p 369.
(17) Cf., for example, R. Scott, D. DePalma, and S. Vinogradov, J. Phys. Chem., 72, 3192 (1968), and references therein.

preliminary studies is in the polar solvent ClCH4-Ticl at 25° with bases 10, 11, 16, 20, 48, and 57 Thies V and VII) indicate that A values are 10 to 20% sy than corresponding values in CCI, except for tri-hylamine which is 10% larger. Thus, with the posble exception of the latter, there is no F nmr evidence of an appreciable enhancement in any hydrogen-bonded on pairs. If the latter were present to a measurable ex-ent in CCl, this form should be further increased in the clar CICH, CH, Cl solvent.

We therefore conclude that in CCl. p-FCiH,OH ornis only the hydrogen-bonded complex with all of the Senes of Table V. Our results are consistent with these lydrogen-bonded complexes having a rather well-deined and a relatively low extent (up to ~30%) of ap-

parent proton transfer.

In accord with the conclusion that there is a small exent of proton transfer in these hydrogen-bonded comblexes is the fact that there is no simple correspondence

(1) (8) Unpublished results of Mr. Joseph Mitzky.

between base strength as measured by the standard free energy of hydrogen-bonded complex formation of p-FC,H,OH in CCl, and corresponding aqueous pK, values of base. Figure 2 shows the complete scatter pattern which results from a plot of pKA 10 vs. log K1 tto values for bases having a variety of common functional groups. This subject is considered in detail in paper V of this series.

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(19) The pK1 values shown in Figure 2 for hazamethylphosphoramide (-1.4) and dimethyl sulfoxide (-2.6) are unpublished results of Rr. J. R. Rukshya, obtained by the method of Haako (P. Haaka, R. D. Cook, and O. H. Hurst, J. Am. Chem. Soc., 89, 2630 (1967)). (20) M. O. Schwartz, Ph.D. Thosis, The Pennsylvania State Uni-

Studies of Hydrogen-Bonded Complex Formation with p-Fluorophenol. V. Linear Free Energy Relationships with OH Reference Acids<sup>1</sup>

versity, Dag 1965.

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Abstract: Linear free energy relationships have been established in the formation of hydrogen-bonded complexes of various OH reference acids with a wide variety of proton acceptors. The effects of temperature, solvent, and substituents have been examined. A unique base parameter, pRes, has been defined which measures the relative strength of the acceptor in hydrogen-bonded complex formation with any suitable OH reference acid. pRine values do not correlate with aqueous pRA values, except within series having a common functional center and variable electronic effects of substituents. pRan values also are not applicable to reference acids involving internal hydrogen bonding and are presumably not applicable to systems in which there is substantial formation of the hydrogen-bonded ion pair (in mobile equilibrium with the hydrogen-bonded complex). Evidence is presented that the pKza scale is applicable (at least qualitatively) to other relatively weak interactions between bases and a "shielded" center of positive charge. The highly dispersed family relationships between page and corresponding pK4 values are indicated to be useful in distinguishing the atomic center of complexing in polyfunctional bases.

The effect of molecular structure on the strength of the hydrogen bond is a subject of diverse intersts. s.a. These include, for example, the structure and cactivity of proteins and many other, natural products, colvation effects in many chemical reactions and in spectroscopic measurements, and transition-state char-Poteristics in proton-transfor processes.

There have been in recent years several extensive tudies of the effects of structure on base strengths

(1) This work was supported in part by the Public Health Service, epicets OM14078 (Irvino), AL-07765 (Princeton), (2) G. C. Pimentel and A. L. McClellan, "The Hydrogen Bond," (2) H. Freeman and Co., San Francisco, Calif., 1960, (3) (a) D. Hedzi, "Hydrogen Bonding," Pergamon Press, New York, J. Y., 1957; (b) W. C. Hamilton and J. A. Iberz, "Hydrogen Bonding in fillds," W. A. Banjamin, Inc., New York, N. Y., 1968, (4) (a) G. Aksnas, and T. Orametad, Acta Chem. Scand., 14, 1475

toward a common reference sold, as measured by the formation constant, K, for the 1:1 hydrogen-bonded

(1960); (b) T. Gramatad, ibid., 15, 1337 (1961); 16, 807 (1962); (c) T. Gramatad and S. I. Saaprud, ibid., 16, 999 (1962); (d) T. Gramatad and W. J. Fuglevik, ibid., 16, 1369, 2368 (1962); (e) T. Gramatad, Specirophim, Acta, 19, 497, 829 (1963); (f) T. Gramatad and W. J. Fuglevik, ibid., 21, 1343 (1965); (g) U. Blindhelm and T. Gramatad, ibid., 21, 1073 (1965); (h) M. D. Joeston and R. S. Drago, J. Am. Chem. Spc., 84, 2696, 3817 (1962); (d) T. D. Episy and R. S. Drago, J. Am. Chem. Spc., 84, 2696, 5817 (1962); (d) T. D. Episy and R. S. Drago, J. Am. Chem. Spc., 85, 1676, 1967); (j) R. West, D. L. Fowell, L. S. Whatley, M. K. T. Lee, and F. von R. Schleyer, ibid., 84, 3221 (1962); (k) D. L. Fowell and R. West, Specirochim, Acta, 20, 983 (1964); (l) R. West, D. L. Powell, M. K. T. Lee, and L. S. Whatley, J. Am. Chem. Soc., 86, 3227 (1964); (m) H. Dunken and H. Fritzicho, Z. Chem., 1, 127 (249 (1961); 1, 345 (1962); (a) Z. Yothlda and B. B. Orawa, J. Am. Chem. Soc., 88, 4019 (1966); (o) C. J. Crawell and A. L. Alired, ibid., 85, 1713 (1963); (b) B. M. Arnett, T. S. S. R. Murty, P. von R. Schloyer, and L. Joris, ibid., B. M. Arnett, T. S. S. R. Murty, P. von R. Schleyer, and L. Joris, ibid., 89, 5955 (1967); (q) D. Gurka, R. W. Taft, L. Joris, and P. von R. Schleyer, ibid., 89, 5957 (1967); (r) D. Gurka and R. W. Taft, ibid., 91,